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<td><strong>Citation</strong></td>
<td>Li, J., Nie, Z., Zheng, Y. Y., Dong, S., &amp; Loh, Z. H. (2013). Elementary electron and ion dynamics in ionized liquid water. The Journal of Physical Chemistry Letters, 4, 3698-3703.</td>
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<td>2013</td>
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Elementary Electron and Ion Dynamics in Ionized Liquid Water

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Abstract

The ionization of liquid water functions as the principal trigger for a myriad of phenomena that are relevant to radiation chemistry and biology. The earliest events that follow the ionization of water, however, remain relatively unknown. Here, femtosecond coherence spectroscopy is combined with polarization anisotropy measurements to elucidate the ultrafast electron and ion dynamics in ionized water. The results show that strong-field ionization of liquid water produces an aligned $p$ electron distribution. Furthermore oscillations observed in the polarization anisotropy are suggestive of valence electron motion in the highly reactive $\text{H}_2\text{O}^+$ radical cation, whose lifetime with respect to proton transfer is found to be $196 \pm 5$ fs. Coherent intermolecular motions that signal initial solvent reorganization and subsequent long-lived ballistic proton transport that involves the $\text{H}_3\text{O}^+$ end-product are also detected in the time domain. These results offer new insight into the elementary dynamics of ionized liquid water.

**Keywords:** Ultrafast spectroscopy, hydrated electron, valence electron motion, hydrogen bond, proton transfer, photoelectron angular distribution
The ionization of liquid water is a universal phenomenon that accompanies the interaction of high-energy radiation with matter in aqueous environments. The ensuing cascade of chemical reactions that involves ions, electrons, and radicals forms the basis of solution and interfacial radiation chemistry.\(^1\) Moreover, since liquid water comprises the major component of cellular matter, oxidative damage to biological systems by high-energy radiation is triggered mainly by the ionization of water.\(^2\) The low-energy electrons that are produced by ionization can also induce radiation damage by dissociative electron attachment to biomolecules.\(^3\) The elementary processes that follow the ionization of liquid water therefore forms the fundamental framework for studies of radiation-matter interaction in chemistry and biology.

The direct products of water ionization are the \(\text{H}_2\text{O}^+\) radical cation and the hydrated electron,\(^1\) both of which are highly reactive species. Ab initio molecular dynamics (AIMD) simulations\(^4,5\) and infrared spectroscopy of ionized water clusters\(^6\) show that \(\text{H}_2\text{O}^+\) undergoes an ion-molecule reaction with a neighboring \(\text{H}_2\text{O}\) molecule to yield the hydronium cation (\(\text{H}_3\text{O}^+\)) and a hydroxyl radical (\(\text{OH}\)). The \(\text{H}_3\text{O}^+\) product subsequently undergoes rapid structural diffusion, akin to the case of an excess proton in water,\(^7\) thereby resulting in the rapid disappearance of the initial \(\text{H}_3\text{O}^+\cdots\text{OH}\) contact pair. The predicted time scale for the proton transfer from \(\text{H}_2\text{O}^+\) is \(\sim 100\) fs,\(^4,5\) although experimental attempts at the determination of its lifetime have thus far been inconclusive.\(^5,8,9\)

In contrast to the elusive \(\text{H}_2\text{O}^+\) cation, the spectroscopy and dynamics of the hydrated electron has been studied extensively.\(^10\) The transition of the hydrated electron from the \(s\)-like ground state to the \(p\)-like excited state underlies its absorption spectrum, which peaks at 719 nm and spans <500 – 1000 nm.\(^11\) Time-resolved optical pump-probe spectroscopy elucidates the ultrafast dynamics following either the injection of an electron into water by multiphoton
ionization (MPI)\textsuperscript{9,12} or the photoexcitation of a pre-equilibrated hydrated electron.\textsuperscript{13,14} The latter studies reveal subpicosecond dynamics with time constants of \~50 fs and \~300 fs, whereby the faster component exhibits a pronounced H/D isotope dependence. The short-time dynamics is ascribed to inertial solvation of the \(p\) state (adiabatic solvation model)\textsuperscript{13} or alternatively, to internal conversion of the \(p\) state to a vibrationally hot \(s\) state (nonadiabatic solvation model);\textsuperscript{14} recent femtosecond photoelectron imaging experiments performed on anionic water clusters lend support to the nonadiabatic solvation model.\textsuperscript{15} Similarly, studies of the hydrated electron generated by MPI reveal the solvation dynamics of the \(s\) state electron, although the intermediary \(p\) state electron has eluded direct spectroscopic observation.\textsuperscript{16–19}

Here, femtosecond coherence spectroscopy\textsuperscript{20–22} is combined with polarization anisotropy measurements to investigate the early-time electron and ion dynamics triggered by the MPI of liquid water. The peak intensity of the 800-nm ionization pump pulse (\(2 \times 10^{13}\) W/cm\(^2\)) yields a Keldysh parameter of \(\gamma \sim 2\) (see Supporting Information). Accordingly the MPI process can be characterized as being in the strong-field regime. Fluence dependence measurements reveal a 9-photon MPI process (see Supporting Information); ionization channels that involve 5- and 6-photon resonance enhancement are accessible at this laser wavelength.\textsuperscript{23} The total energy deposition of 14.0 eV is significantly above the vertical ionization potential of liquid water (11.16 eV).\textsuperscript{24} At this input energy, the large electron ejection length of \(\langle r_0 \rangle \sim 35\) Å,\textsuperscript{25} which defines the spatial extent of the surrounding medium over which the electron is delocalized, allows access to the collective dynamics of up to \(\sim 10^4\) water molecules. The 800-nm probe pulse interrogates the \(s \rightarrow p\) transitions of both the hydrated \(s\) electron and its pre-solvated precursor.\textsuperscript{26,18} The manner in which the intermolecular vibrational coherences of liquid water modulates the electron \(s \rightarrow p\) absorption is analogous to the way in which a vibrational wave packet that is
Figure 1. Femtosecond time-resolved differential absorption signal ($\Delta$OD) as a function of time delay for parallel ($S_\parallel$) and perpendicular ($S_\perp$) relative polarizations between pump and probe beams. The inset shows the $S_\parallel$ signal at later time delays, during which an oscillation is apparent.

launched on a molecular potential energy surface modifies the energy and amplitude of its associated electronic transitions.\textsuperscript{27,28} By relying on the observation of coherences and polarization anisotropies that are associated with the various intermediates, our approach complements earlier methods based on optical pump-probe transient absorption (TA) spectroscopy, which has its limitations. For example, the inability to isolate the H$_2$O$^+$ species in a recent femtosecond deep UV pump-supercontinuum probe TA experiment was attributed to its weak absorption being overwhelmed by the strong absorbance of the solvated electron.\textsuperscript{5}

The time-dependent differential absorption time traces for parallel ($S_\parallel$) and perpendicular ($S_\perp$) relative pump-probe polarizations are shown in Figure 1. Beyond the region of pump-probe temporal overlap (~50 to +50 fs), where nonlinear effects that involve both pump and probe beams dominate, both $S_\parallel$ and $S_\perp$ are positive and show an overall increase with time delay that can be attributed to the formation of the hydrated $s$ electron. The ionization fraction is estimated to be $7 \times 10^{-6}$ based on the differential absorbance measured at 1-ps time delay and the known molar extinction coefficient of the hydrated electron at 800 nm (1.6 \times 10^4 M^{-1}cm^{-1}).\textsuperscript{26} Note that the recombination of the hydrated electron with OH and/or H$_3$O$^+$ do not contribute significantly
Figure 2. (a) The polarization anisotropy $r$ shows that the $p$ electron is produced by the MPI of liquid water. The symbols correspond to the experimental data. The solid lines are obtained from the fits to the theoretical model (see main text). (b) Time-evolution of the $p$ state electron angular distribution in liquid H$_2$O, as reconstructed from experimental data.

The polarization anisotropy observed at early times provides direct evidence for the creation of an aligned electron by the MPI of liquid water (Figure 2a). The time-dependent polarization anisotropy is modeled by considering that strong-field ionization by a laser pulse with electric field polarized along the $z$ direction produces an incoherent mixture of $s$ and $p_z$ electrons with initial fractional populations $f_s(0)$ and $f_{p_z}(0)$, respectively (see Supporting Information). The model further assumes that the $p_z$ excited state relaxes to the $s$ ground state...
with a decay lifetime $\tau_p$. While electron states with higher angular momenta can be populated by ionization, the angular momenta states that are included in our model are restricted to the $s$ and $p$ states, due to the predominant sensitivity of the 800-nm probe wavelength to the $s \rightarrow p$ transition, as well as the limitations on the number of independent variables that can be extracted from the data. Moreover omission of the $p_x$ and $p_y$ electrons from the above consideration is motivated by previous work on gas-phase atomic and molecular strong-field MPI, which show that angular distributions of the ejected electrons are primarily aligned along the polarization axis of the ionization pump pulse.\textsuperscript{37,38} Fitting of the polarization anisotropy with only $f_p(0)$ and $\tau_p$ as adjustable parameters yields, in the case of $\text{H}_2\text{O}$ ($\text{D}_2\text{O}$), values of $0.94 \pm 0.05$ ($0.92 \pm 0.04$) for $f_p(0)$ and $79 \pm 5$ fs ($101 \pm 6$ fs) for $\tau_p$ (Figure 2a). The reconstructed temporal evolution of the electron angular distribution following the ionization of liquid $\text{H}_2\text{O}$ is shown in Figure 2b.

The creation of an aligned electron distribution in this work is facilitated by the use of a strong-field-ionizing pump pulse, whereas it should be noted that previous MPI studies of liquid water were performed with pump pulse intensities that are in the perturbative regime. While the initial fractional populations of the $p_z$ electron generated in $\text{H}_2\text{O}$ and $\text{D}_2\text{O}$ are identical to within experimental error, the retrieved $p$ state lifetimes are clearly dependent on H/D isotopic substitution, with the lifetime being longer in $\text{D}_2\text{O}$ than in $\text{H}_2\text{O}$ by a factor of $1.3 \pm 0.1$; note that a more negative anisotropy at 60-fs time delay in $\text{D}_2\text{O}$ than in $\text{H}_2\text{O}$ is consistent with the longer $p$ electron lifetime in the former (Figure 2a). This result is reminiscent of that obtained from previous studies of the $p$ electron dynamics that follows the photoexcitation of the pre-equilibrated $s$ electron.\textsuperscript{14,15} In those studies, the factor-of-$\sqrt{2}$ slowdown observed in the internal conversion dynamics upon deuteration is attributed to the participation of librational modes in electronic relaxation. In fact, we note the similarity in relaxation time scales for the $p$ electron
prepared by MPI and that obtained within the nonadiabatic solvation model of photoexcited hydrated electron dynamics. The H/D substitution dependence observed here therefore suggests that solvent librational modes mediate the relaxation of the injected $p$ state electron prior to its solvation. The slightly longer lifetime of $p$ electrons produced by MPI than by photoexcitation can be rationalized in terms of differing initial conditions: the solvation of an electron following the ionization of water is likely to involve a larger solvent reorganization penalty than the solvation of a partially hydrated $p$ electron produced by photoexcitation.

Oscillatory features in the polarization anisotropy decay indicate a periodic reorientation of the probe transition dipole moment that is concomitant with the electronic relaxation of the $p$ state electron. The residual polarization anisotropies, obtained by subtracting the fits for the $p$ electron population decay from the polarization anisotropy signals over a time delay range of 60 – 350 fs, are shown in Figure 3a for both H$_2$O and D$_2$O. Their corresponding Fast Fourier Transform (FFT) power spectra reveal oscillation frequencies of 150 cm$^{-1}$ and 500 cm$^{-1}$ (Figure 3b, solid lines), neither of which exhibits a perceptible shift with H/D substitution. Time-domain analysis yields, for the case of H$_2$O (D$_2$O), cosinusoidal phases (in units of $\pi$ rad) of $-0.31 \pm 0.09$ ($-0.39 \pm 0.09$) and $1.30 \pm 0.08$ ($1.00 \pm 0.04$) for the low- and high-frequency component, respectively (see Supporting Information). The exponential damping times of both frequency components in the oscillations are $70 \pm 12$ fs and $64 \pm 7$ fs for H$_2$O and D$_2$O, respectively. For comparison, the FFT power spectra of the isotropic signals $S_{iso}$, where $S_{iso} = (S_\parallel + 2S_\perp)/3$ (Figure 3b, dashed lines), exhibit only one frequency component at 150 cm$^{-1}$.

A possible source of the 500-cm$^{-1}$ polarization anisotropy modulation is solvent librational motion. The sub-100-fs lifetime of librational excitation, as elucidated by ultrafast infrared spectroscopy and molecular dynamics simulations, is consistent with the $\sim$70-fs
Figure 3. (a) The residual polarization anisotropy (symbols) reveals oscillatory features, which can be fit to a sum of two exponentially decaying cosinusoidal functions (solid lines). (b) The FFT power spectra of the residual polarization anisotropies reveal frequency components at 150 and 500 cm\(^{-1}\) (solid lines), both of which are independent of H/D substitution. For comparison, the FFT power spectra of the isotropic signals are also shown (dashed lines).

decay of the polarization anisotropy oscillation observed here. Despite the broad FFT frequency linewidth, however, it is unclear as to why the 500-cm\(^{-1}\) component is centered on the L\(_1\) librational band of H\(_2\)O, even though the L\(_2\) band at 780 cm\(^{-1}\), which extends to ~1000 cm\(^{-1}\), bears considerable Raman intensity.\(^{41}\) Furthermore the 500-cm\(^{-1}\) band does not shift with H/D isotope substitution, even though the librational frequencies of D\(_2\)O are lower than those of H\(_2\)O by \(\sqrt{2}\times\).

In addition to librational motion, another potential explanation for the 500-cm\(^{-1}\) component is angular reorientation via electron motion. Indeed, previous studies of multichromophoric systems have shown that coherent electronic energy transfer leads to
Figure 4. Orbital densities of the (a) HOMO, (b) HOMO–1, and (c) HOMO–2 of (H₂O)₄. Note the orthogonal directions of the dominant orbital densities of the different molecular orbitals, which suggests that their coherent superposition would give rise to a periodic reorientation of the hole density in the (H₂O)₄⁺ ion.

oscillations in the polarization anisotropy signal that are absent in the isotropic signal. It has recently been shown that the strong-field MPI of atoms can trigger valence electron motion in the resultant ions, for which the ~1-eV energy spacing $\Delta E$ between valence electronic states leads to few-femtosecond oscillation periods $T_{el} = \hbar/\Delta E$. In the case of liquid water, the initiation of valence electron motion by strong-field MPI is favored by its dense manifold of electronic states, which presents aqueous H₂O⁺ ion states with energetic separations that are commensurate with the 500-cm⁻¹ oscillation frequency observed here. Furthermore a recent theoretical study suggests that the sudden ionization of the water dimer induces the oscillation of charge density between the oxygen centers of the (H₂O)₂⁺ cation. Our ab initio calculations on the (H₂O)₄ model cluster suggests, within a Koopman-type picture, the common occurrence of (H₂O)₄⁺ ion states with orthogonal hole densities. For example, the orbital densities of the HOMO, HOMO–1, and HOMO–2 orbitals of (H₂O)₄, which correspond to the hole densities of the three lowest (H₂O)₄⁺ ion states, are each oriented in perpendicular directions (Figure 4). The coherent superpositions of these ion states leads to a periodic reorientation of the hole density, which in turn modulates the alignment of the $p$ electron via the Coulomb interaction, from which
an oscillatory component in the polarization anisotropy results. The coherent electronic origin of
the observed 500-cm\(^{-1}\) oscillation is further supported by its cosinusoidal phase of \(\sim \pi\) rad, which
implies a polarization anisotropy that begins at its negative extremum. That is, the hole density at
the moment of ionization is maximally aligned, as observed previously for coherent atomic ion
states prepared by strong-field MPI.\(^{44}\) It is important to note here that the \(\sim 70\)-fs dephasing time
of the hole coherence is not limited by the 1.6-fs optical dephasing time of the \(s \rightarrow p\) transition of
the hydrated electron.\(^{48,49}\)

The 150-cm\(^{-1}\) component that arises in both isotropic and polarization anisotropy signals
is assigned to a Raman-active intermolecular hindered translation along the hydrogen bond
coordinate.\(^{41}\) From previous time-domain optical Kerr effect studies of pure water\(^{50}\) and aqueous
salt solutions,\(^{51}\) the emergence of this mode in \(S_{iso}\), despite its near-complete Raman
depolarization for pure liquid water,\(^{41}\) is a manifestation of the solvent reorganization that
follows the MPI of water. This conclusion is further supported by resonance Raman
measurements\(^{52}\) and theoretical calculations\(^{53,54}\) on the hydrated \(s\) electron, which show that its
photoexcitation is coupled to the hindered translation of solvent molecules. On the other hand,
the appearance of the 150-cm\(^{-1}\) component in the polarization anisotropy is ascribed to hindered
intermolecular translation that is driven by the periodic reorientation of the hole density. In
support of this assignment, an ab initio calculation of the \((H_2O)_4^+\) radical cation reveals hindered
translational modes with frequencies in the range of 140 – 210 cm\(^{-1}\) and depolarization ratios of
0.4 – 0.6 (see Supporting Information). The observed phase shift between the 500-cm\(^{-1}\) and 150-
\(\text{cm}^{-1}\) frequency components can be rationalized in terms of the nonadiabatic driving of the
hindered translational mode by the electron motion.
Figure 5. (a) The time-frequency spectrogram for the anisotropic signal shows the disappearance of the 150 and 500 cm\(^{-1}\) frequency components at early times and the concomitant rise of the frequency component at 310 cm\(^{-1}\). (b) The FFT amplitude at 310 cm\(^{-1}\) reveals an oscillatory rise (symbol), from which a fit to first-order kinetics (solid line) yields a lifetime of 196 ± 5 fs for the aqueous H\(_2\)O\(^+\) radical cation.

The time-frequency spectrogram derived from the anisotropic signal \(S_{aniso} = S_\parallel - S_\perp\) (Figure 5) shows the decay of frequency components at 150 cm\(^{-1}\) and 500 cm\(^{-1}\) and the appearance of a 310-cm\(^{-1}\) component. The new frequency component originates from the oscillatory feature that extends to 1 ps in both \(S_\parallel\) and \(S_\perp\) signals (Figure 1 inset). Note that the center frequency of the nominal 310-cm\(^{-1}\) component shifts periodically in the range of 260 – 330 cm\(^{-1}\); the period of the shift corresponds to a frequency of 60 cm\(^{-1}\).

The feature at 310 cm\(^{-1}\), which is independent of H/D substitution, is attributed to the hindered translation between water molecules and the H\(_3\)O\(^+\) hydronium cation to which they are bound, \textit{cf.} 175 cm\(^{-1}\) for pure water.\(^{41}\) The vibrational frequency blueshift upon water protonation,
which implies the strengthening of the hydrogen bond, is consistent with the contraction of the intermolecular O⋯O distance from 2.85 Å in pure water to 2.52 Å in aqueous acid. This assignment is further supported by ab initio results for the Zundel ion (H₉O₄⁺), which reveal hindered translational modes involving the central H₃O⁺ moiety and the peripheral H₂O molecules at 270 cm⁻¹ and 330 cm⁻¹ (see Supporting Information). The latter, being the asymmetric combination, is completely depolarized and is therefore expected to feature prominently in the anisotropic signal. The 60-cm⁻¹ modulation of the hindered translation mode suggests that this mode is coupled to the hydrogen bond bending vibration.

The extended appearance of the 310-cm⁻¹ frequency component in the spectrogram signifies long-lived coherent intermolecular hindered translational motion that accompanies ballistic proton transport. This vibrational coherence is reaction-driven, in this case, by the strongly exergonic initial proton transfer from H₂O⁺ to a neighboring water molecule. AIMD studies show that this intermolecular vibrational mode drives excess proton transport in water: the proton donor and proton acceptor water molecules must first approach one another before proton transfer is energetically feasible. Moreover, the observed ballistic proton transport agrees with the non-diffusive nature of proton transfer predicted by AIMD simulations of ionized liquid water.

Since H₃O⁺ is initially produced by the transfer of a proton from the H₂O⁺ radical cation to a neighboring water molecule, the onset of ballistic proton transport is expected to correlate with the population decay of the H₂O⁺ species. Fitting the growth of the 310-cm⁻¹ frequency component to pseudo-first-order kinetics yields a lifetime of 196 ± 5 fs for the H₂O⁺ radical cation. This experimentally determined lifetime is significantly longer than the ~100-fs lifetime predicted by AIMD simulations. However we note that the simulations did not consider the
presence of a hydrated electron, or the possibility that aqueous H$_2$O$^+$ could exist as a coherent superposition of delocalized ion states. It is possible that solvent reorganization to accommodate the injected electron competes with the solvation of H$_2$O$^+$, thereby impeding proton transfer to produce H$_3$O$^+$. Moreover, the H$_2$O$^+$$\cdots$H$_2$O bond contraction that is necessary for proton transfer$^5$ from H$_2$O$^+$ occurs on a time scale that is $\sim$3$\times$ slower than the coherent charge oscillation of the aqueous H$_2$O$^+$ hole density. The altering identity of the proton donor conceivably frustrates the proton transfer step, thus prolonging the lifetime of the H$_2$O$^+$ radical cation.

The present work offers a glimpse of the multitude of hitherto unobserved elementary electron and ion dynamics that are triggered by the ionization of liquid water. These experimental results provide strong impetus for the development of a comprehensive theoretical treatment of liquid water ionization that considers the coupled dynamics of the injected electron, the H$_2$O$^+$ radical cation, and the solvent medium. Furthermore, it is envisioned that experiments with improved time resolution will uncover coherent dynamics that evolve on shorter time scales, such as those triggered by core-level ionization,$^{60}$ or those related to intramolecular vibrational motion and/or electron motion involving different electronic bands. Finally, we emphasize that the coherent dynamics elucidated in this work remain pertinent to the study of water ionization by high-energy charged particles, since the rapid traversal of a charged particle through a molecule is equivalent to exposing the molecule to an ultrashort electromagnetic pulse.$^{61}$ With typical traversal times that easily approach the sub-femtosecond regime,$^{62}$ such high-energy charged particles could certainly trigger a vast array of coherent dynamics beyond those reported here.
Acknowledgments

We are greatly indebted to Prof. M. Parrinello and Dr. A. Hassanali for illuminating discussions. We also thank Prof. H.-S. Tan for an equipment loan, and Prof. C. Andreani, Prof. R. Santra and Dr. O. Vendrell for useful discussions. This work is supported by a start-up grant from NTU, the A*Star Science and Engineering Research Council Public Sector Funding (122-PSF-0011), and the award of a Nanyang Assistant Professorship to Z.-H.L.

Supporting Information

Experimental and computational methods, Cartesian coordinates of the optimized geometries of (H₂O)₄ and H₉O₄⁺ obtained from ab initio calculations, calculated vibrational frequencies, Raman activities, and depolarization ratios of (H₂O)₄⁺ and H₉O₄⁺, results of fluence dependence measurements for the determination of the pump photon order, derivation of the theoretical model for fitting the polarization anisotropy decay, time-domain analysis of the residual polarization anisotropy decay, and calculation of the frequency-time spectrogram and its application to obtaining the lifetime of aqueous H₂O⁺. This material is available free of charge via the Internet at http://pubs.acs.org.
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